[1,2,4]TRIAZOLO[1,5-c]PYRIMIDINE DERIVATIVES

Publication number: WO03068776

Publication date:

2003-08-21

Inventor:

IMMA HIRONORI (JP); WATANABE TOMOKAZU (JP);

SHIOZAKI SHIZUO (JP); KANDA TOMOYUKI (JP); KUWANA YOSHIHISA (JP); SHIMADA JUNICHI (JP)

Applicant:

KYOWA HAKKO KOGYO KK (JP); IMMA HIRONORI (JP); WATANABE TOMOKAZÙ (JP); SHIOZAKI SHIZUO (JP); KANDA TOMOYUKI (JP); KUWANA

YOSHIHISA (JP); SHIMADA JUNICHI (JP)

Classification:

- international: A61P3/10: A61P9/00: A61P9/10: A61P25/16;

A61P25/24; A61P25/28; A61P43/00; C07D487/04; C07D519/00; A61P3/00; A61P9/00; A61P25/00; A61P43/00; C07D487/00; C07D519/00; (IPC1-7):

C07D487/04; A61K31/519; A61K31/5377; A61K31/5383;

A61K31/542; A61P3/10; A61P9/00; A61P9/10; A61P25/16; A61P25/24; A61P25/28; A61P43/00;

C07D519/00

- European: C07D487/04; C07D519/00 Application number: WO2003JP01565 20030214 Priority number(s): JP20020037819 20020215

Also published as:

🔯 AU2003211993 (A1)

Cited documents:



WO9842711 WO0017201

Report a data error here

Abstract of WO03068776

[1,2,4]Triazolo[1,5-c]pyrimidine derivatives represented by the following general formula (I) or pharmacologically acceptable salts thereof which show an adenosine A2A receptor antagonism and are useful in treating and/or preventing various diseases caused by the hyperfunction of the adenosine A2A receptor: (I) wherein R1 represents optionally substituted anyl or optionally substituted heteroaryl; R2 represents hydrogen, halogeno, optionally substituted lower alkyl, optionally substituted aryl or optionally substituted heteroaryl; and R3 represents a group of any of the following formulae (A), (B) and (C).

Data supplied from the esp@cenet database - Worldwide